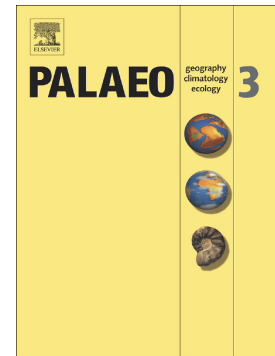


Accepted Manuscript

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PII: S0031-0182(19)30141-5

DOI: <https://doi.org/10.1016/j.palaeo.2019.04.009>

Reference: PALAEO 9153

To appear in: *Palaeogeography, Palaeoclimatology, Palaeoecology*

Received date: 11 February 2019

Revised date: 5 April 2019

Accepted date: 8 April 2019

Please cite this article as: S.R. Franch, J. Echevarría, S.E. Damborenea, et al., Population response during an Oceanic Anoxic Event: The case of Posidonotis (Bivalvia) from the lower Jurassic of the Neuquén Basin, Argentina, *Palaeogeography, Palaeoclimatology, Palaeoecology*, <https://doi.org/10.1016/j.palaeo.2019.04.009>

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**Population response during an Oceanic Anoxic Event: the case of
Posidonotis (Bivalvia) from the Lower Jurassic of the Neuquén Basin,
Argentina**

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Abstract. Benthonic marine species show a wide range of biological reactions to seawater chemical changes through time, from subtle adjustments to extinction. The Early Toarcian Oceanic Anoxic Event (T-OAE) was recently recognized in the Neuquén Basin, Argentina, confirming its global scope. The event was identified

chemostratigraphically on the basis of a relative increase in marine organic carbon and a characteristic negative carbon-isotope excursion ($\delta^{13}\text{C}_{\text{org}}$) in bulk rock and fossil wood in the upper Pliensbachian–lower Toarcian interval in the Arroyo Lapa section (Neuquén). Simultaneously with collection of lithological samples, a high-resolution biostratigraphical survey was carried out, and the scarce benthonic fauna was collected in order to check the biotic response to changing marine geochemical conditions. We present here an analysis of size and abundance data from the T-OAE interval in the Neuquén Basin for the dominant bivalve species, the paper-clam *Posidonotis cancellata* (Leanza), and relate these data to geochemical proxies (%TOC and $\delta^{13}\text{C}_{\text{org}}$) obtained at the same locality. The abundance of *P. cancellata* increased when the rest of the benthos diminished, reaching a maximum at the onset level of the T-OAE, and then decreasing. Size-frequency distributions show a noteworthy lack of juvenile shells. Shell size shows a positive correlation with %TOC in the whole section, though over the T-OAE interval proper, it decreases below the level where the maximum %TOC value is attained and increases above it. *Posidonotis cancellata* shows features of opportunistic species, such as high tolerance to hypoxia, strong dominance in impoverished environments and a strong dependence on primary productivity, but at the same time had a reproductive strategy more similar to equilibrium species, with relatively low juvenile mortality rates. Several anatomical features suggest adaptation to permanently dysaerobic environments. The species disappeared just before the minimum negative carbon-isotope value was reached; and by the same time the genus became extinct worldwide.

Key words: Toarcian OAE, opportunistic species, biotic reactions, paper-clams, South America.

1. Introduction

A relatively high extinction rate affecting various sorts of organisms was recorded around the world for the late Pliensbachian–early Toarcian interval, with a peak at the *Tenuicostatum/Falciferum* ammonite zonal boundary (Raup and Sepkoski, 1984; Hallam, 1986; Sepkoski and Raup, 1986; Little and Benton, 1995; Hallam and Wignall, 1997; Harries and Little, 1999; Aberhan and Fürsich, 2000; Macchioni and Cecca, 2002; Cecca and Macchioni, 2004; Boomer et al., 2008; Caswell et al., 2009; Dera et al., 2010; Guex et al., 2012; Fantasia et al., 2018). This early Toarcian extinction event has been linked to the Early Toarcian Oceanic Anoxic Event (T-OAE) (Hallam, 1986, 1987; Aberhan and Baumiller, 2003; Wignall et al., 2005; Gómez et al., 2008; Bond and Wignall, 2014) and/or to global warming (Bailey et al., 2003; Dera et al., 2009; Dera and Donadieu, 2012; García Joral et al., 2011; Gómez and Goy, 2011; Danise et al., 2013; Krencker et al., 2015; Dunhill et al., 2018). The T-OAE was characterized by widespread deposition of shales rich in organic carbon, a negative carbon-isotope excursion (CIE) typically intersecting an overall positive shift, local euxinic conditions and seawater temperature rise (Jenkyns, 1985, 1988, 2003, 2010; Jenkyns and Clayton, 1986, 1997; Hesselbo et al., 2000, 2007; Hermoso et al., 2009; Danise et al., 2013; Xu et al., 2018), and is regarded as one of the most significant environmental perturbations of the Mesozoic Era.

The ecological consequences of this particular extinction event are relevant to understanding how all the inter-connected environmental changes may have affected marine organisms (Röhl et al., 2001; Caswell et al., 2009; Morten and Twitchett, 2009; Dera et al., 2010; Fraguas and Young, 2011; García Joral et al., 2011; Danise et al., 2013; Caswell and Frid, 2013, 2016; Caswell and Coe, 2014; Danise et al., 2019).

Abundance and body size are two ecologically related parameters that have been studied in this context. It has been suggested that body size was related to extinction dynamics in different organisms (McKinney, 1990, 1997; Jablonski, 1996; Payne, 2005; Twitchett, 2007; Harries and Knorr, 2009; Piazza et al., 2019). In this context, there are a few analyses of the effects on size and other morphological features of marine invertebrates and vertebrates during the early Toarcian extinction event (Mortén and Twitchett, 2009; Maxwell and Vincent, 2015; García Joral et al., 2018). For example, detailed quantitative studies on abundance and size on populations of the bivalves *Bositra radiata* (Goldfuss) and *Pseudomytiloides dubius* (Sowerby) from T-OAE black shales in Yorkshire, UK, in relation to geochemical proxies, yielded insightful results (Caswell and Coe, 2013) that can be compared with those presented here.

The genus *Posidonotis* Losacco, 1942 (including its junior synonym *Pectinula* Leanza, 1943) has a stratigraphic range restricted to the Lower Jurassic (upper Sinemurian to lower Toarcian). Its distribution was reviewed by Damborenea (1989) and Aberhan and Pálffy (1996): in North America, for example, the genus occurs in upper Sinemurian to lower Toarcian beds, whilst in South America and elsewhere it is restricted to upper Pliensbachian to lower Toarcian deposits (Fig. 1). Although most abundant in temperate regions of the eastern Palaeo-Pacific, it has also sporadic occurrences in central Tethys (Italy: Losacco, 1942; Monari, 1994; Greece: Damborenea, 1987), and Japan (Hayami, 1988; Tanabe, 1991), but it appears to be absent from very high palaeolatitudes. The Tethyan and Japanese records were attributed by Aberhan and Pálffy (1996) to eastward migration along the Hispanic Corridor and subsequent isolation. Although this was the most probable migration route for benthonic bivalves (Damborenea and Manceñido, 1979), *Posidonotis* distribution data may also be explained by dispersion across the Pacific following the westward

direction of prevailing currents (Damborenea et al., 2013, fig. 6.4). Throughout its geographical and temporal range, *Posidonotis* commonly occurs forming monospecific shell pavements in dark/black shales. The functional morphology, facies distribution and palaeoecology of *Posidonotis* species were extensively discussed by Damborenea (1987, 1989) and Aberhan and Pálffy (1996), who regarded them as opportunistic species characteristic of quiet-water, dysaerobic environments.

The species *Posidonotis cancellata* (Leanza) was systematically revised by Damborenea (1987) and was then provisionally regarded as a distinct species on the basis of subtle morphological differences. On the other hand, Aberhan and Pálffy (1996) considered it as a subjective synonym of the long-ranging *P. semiplicata* (Hyatt). In Argentina, the species has a very wide geographical distribution from San Juan Province in the north to Chubut Province in the south (location of records from Mendoza and Neuquén provinces are shown in Fig. 1B), although it is restricted stratigraphically to the uppermost Pliensbachian–lowermost Toarcian interval (*Fanninoceras disciforme* to *Tenuicostatum* Zones). The type-species of the genus, *Posidonotis dainelli* Lossaco was present in the Mediterranean (Greece and Italy) Tethys (Losacco, 1942; Damborenea, 1987, 1989; Monari, 1994) and Japan (Hayami, 1988; Tanabe, 1991) in comparable sedimentary environments at about the same time (Izumi et al., 2012).

Recently, the sedimentary record of the T-OAE was recognized in the Neuquén Basin, Argentina (Fig. 1), confirming the event as a global phenomenon (Al-Suwaidi et al., 2010, 2016). The event was identified chemostratigraphically on the basis of a relative increase in marine organic carbon and a characteristic overall positive excursion, intersected by a negative carbon-isotope excursion ($\delta^{13}\text{C}_{\text{org}}$) in bulk rock and fossil wood. High-resolution sampling with tight biostratigraphical control (ammonites,

Riccardi, 2008a, 2008b) was carried out in the upper Pliensbachian/lower Toarcian interval in the Arroyo Lapa section (Fig. 2), where benthonic species are scarce.

Although other authors have suggested a general correlation between degree of oxygenation and shell size for *Posidonotis* species (e.g., Aberhan and Pálffy, 1996) no statistical analysis had been attempted prior to this study. Hence, we here present a thorough analysis of size and abundance data from the T-OAE interval in the Neuquén Basin for the dominant bivalve species, the paper-clam *Posidonotis cancellata* (Leanza), and relate these data to geochemical proxies [total organic carbon (%TOC) and $\delta^{13}\text{C}_{\text{org}}$].

2. Geological setting

The Neuquén Basin is an extensive Mesozoic back-arc basin developed on the western convergent margin of the South American plate (Legarreta and Uliana, 1996; see also Fig. 1), in connection with the break-up of Gondwana (Uliana and Biddle, 1988). After initial rifting (spanning the Late Triassic to earliest Jurassic), during the sag phase (Sinemurian to Toarcian) the coalescence of formerly unconnected half-grabens took place with widespread transgression over most of the Neuquén embayment (Vergani et al., 1995; Legarreta and Uliana, 1996).

The presence of a well-developed Lower Jurassic succession at Arroyo Lapa (Charahuilla-Chacaico area) was recognized long ago (e.g., Weaver, 1931, p. 27-29; Groeber et al., 1953, p. 158-161; Volkheimer, 1973). There, the Lower–Middle Jurassic Cuyo Group attains about 1500 m in thickness. From the lithostratigraphical point of view, the interval considered in this study is consistently assigned to the Los Molles Formation (Weaver, 1931; see also Riccardi and Damborenea, 1993; Gulisano and Gutiérrez-Pleimling, 1995; Arregui et al., 2011). Furthermore, paleomagnetic data obtained from this section were used to build a composite magnetostratigraphy for the

Early Jurassic of the Neuquén Basin (Iglesia Llanos and Riccardi, 2000). Riccardi (2005) and Manceñido et al. (2007) first reported the occurrence of organic-rich dark shales probably representing the T-OAE in this section, which was subsequently confirmed with detailed sedimentological and geochemical evidence by Al-Suwaidi et al. (2010, 2016, see above).

The surveyed section of the Los Molles Formation (Fig. 2A) starts with conglomerates overlying tuffs of the Lapa Formation, followed by about 80 m of lower Pliensbachian deposits dominated by sandstones with interbedded siltstones. Upper Pliensbachian deposits comprise about 150 m of heterogeneous siliciclastic and pyroclastic sediments of alternating sandstones, siltstones and tuffs. The next package, over 70 m thick, spans the latest Pliensbachian to earliest Toarcian time interval, and consists of laminated black shales and siltstones interbedded with relatively thin sandstone layers (Fig. 2B, further description in Al-Suwaidi et al., 2010). This last interval was logged and sampled in detail and provided the data for this study.

3. Material and methods

3.1. Material

The analysis performed here was restricted to material from the Arroyo Lapa South section, because detailed geochemical data (published by Al-Suwaidi et al., 2010) were obtained from rock samples taken simultaneously as the paleontological samples (Ros-Franch et al., 2014). Albeit additional detailed data for the T-OAE interval are available (Al-Suwaidi et al., 2016), they were not used here in order to maintain statistical consistency, since they only cover a minor (upper) part of the stratigraphic range of *P. cancellata* in the logged section. The fossil material is deposited in the Invertebrate Palaeontology collection of the La Plata Museum (MLP 35869 to 35903).

Size data were obtained from 502 photographed individual valves. Shell length and surface area (Fig. 3) were measured using ImageJ software (<http://rsb.info.nih.gov/ij/>). A synthesis of the data used is shown in Table 1. Abundance data were recorded as: (a) area covered by specimens on each slab/slab surface area (Fig. 4); (b) number of specimens present per dm^2 of each slab surface.

3.2. Variables analyzed

The main goal of the present paper is to check for responses in *Posidonotis* populations associated to variations in total organic carbon (%TOC) and carbon-isotope ($\delta^{13}\text{C}_{\text{org}}$) ratios. %TOC can potentially indicate changes in primary production, assuming that this quantity factor is the first-order control on organic-matter flux to the sea floor and subsequent burial, rather than variable preservation (cf. Demaison and Moore, 1980; Pedersen and Calvert, 1990). Variations in bulk $\delta^{13}\text{C}$, if recording global changes in the ocean–atmosphere system, should relate to ρCO_2 , which will be drawn down during episodes of global organic-matter burial such as the T-OAE, if not counterbalanced by methanogenic and/or volcanic sources, to which the characteristic negative excursion is usually ascribed (Hesselbo et al., 2000; McElwain et al., 2005; Hermoso et al., 2012; Trecalli et al., 2012; Ullmann et al., 2014). Evidence for volcanism, which would have supplied CO_2 to the atmosphere, is given by the stratigraphic association of relatively elevated contents of mercury within sediments deposited during the T-OAE in many places, including Arroyo Lapa South, with an ultimate source suggested as the Karoo–Ferrar Large Igneous Province (Percival et al., 2015, 2016).

When checking for association between population and environmental variables, two options were considered: direct correlation to the variable (comparison with %TOC

and $\delta^{13}\text{C}_{\text{org}}$ closest to the *Posidonotis* sample) and correlation to trends on the variables (first %TOC and $\delta^{13}\text{C}_{\text{org}}$ value after the *Posidonotis* sample minus immediate previous values, Table 1, columns "dif. %TOC" and "dif. $\delta^{13}\text{C}_{\text{org}}$ "; when the *Posidonotis* sample coincided with a geochemical sample, the trend was determined as sample value minus the immediate previous value).

Variables analyzed for *Posidonotis* samples (Table 1) were individual sizes distribution and population density. Only samples with at least five measured shells were considered. Maximum size was quantified as maximum-length per sample, and five maximum-length values per sample. Maximum-length per sample is often used as a size estimator (e.g., Hallam, 1975; Aberhan and Fürsich, 1997; Morten and Twitchett, 2009), but it can be strongly influenced by outliers. Hence, the analyses were also performed on the five largest measurements per sample. Nevertheless, these two data sets seem to be dependent on sample size (for maximum length Spearman's $r_s = 0.53$, $p = 0.022$; for 5 maximum lengths $r_s = 0.52$, $p < 0.01$); *i.e.*, the larger the sample, the higher the probability of finding large shells.

For length-data distribution, the medians, variances, skewness and kurtosis were also considered. Anderson (2001) used size variance as an estimate of size range, but also as a proxy for maximum size. Since the lowest value is bounded (values less than zero being impossible), variance will increase primarily due to larger values. Variance will also be less sensitive to outliers than maximum length. Levene's test for homogeneity of variances shows significant differences either based on means or medians ($p < 0.01$), and there seems to be no correlation between sample size and variance ($r_s = 0.38$, $p = 0.12$).

3.3. Methodology

The geochemical variables and the size and abundance data are expected to be responding to the same environmental factors (those responsible for the OAE); hence a monotonic (not necessarily linear) association should be expected among these sets of variables. The non-parametric Spearman rank-order correlation coefficient (r_s) was used to check that association. A correlation analysis was also performed for size and abundance data with reference to stratigraphic position during the T-OAE portion of the section, to check for trends over time.

From the differences within the geochemical variables (Table 1, columns "dif. %TOC" and "dif. $\delta^{13}\text{C}_{\text{org}}$ "), another analysis was proposed in order to compare with the results provided by Caswell and Coe (2013) for *Bositra radiata*. The samples were grouped in positive and negative differences for both geochemical variables (treated as increasing and decreasing trends), and the length-frequency distributions for both groups were compared by means of a Kolmogorov–Smirnov test (to check for equality of distribution) and Mann–Whitney U test (to check for equality of medians). All analytical methods are described in Hammer and Harper (2006) and were performed on PAST 3.20 (Hammer et al., 2001).

4. Results

The typical matrix of *Posidonotis cancellata* consists of fairly thin-layered, dark shales, bearing monospecific pavements, made up of flat-lying, concordant valves (Fig. 2C). The valves are fragile, thin-shelled, usually unbroken (even if disarticulated), which suggests undisturbed, autochthonous accumulation. Most commonly, specimens appear as internal and external moulds (Fig. 4B), although in some instances delicate shell remains are preserved as white paper-thin laminae (Fig. 4C).

Size and abundance data show a general correspondence (Fig. 5), though only the 5 maximum-length values show a significant negative response to values of, and changes in, $\delta^{13}\text{C}_{\text{org}}$ (Table 2). These results seem to be dependent on sample size.

Skewness, on the other hand, seems to be positively correlated with %TOC values.

Comparison of groups of samples under increasing and decreasing trend contexts in geochemical variables (Fig. 6) shows a significant difference in distribution and median for both %TOC (Kolmogorov–Smirnov test $p < 0.01$; Mann-Whitney U test $p < 0.01$) and $\delta^{13}\text{C}_{\text{org}}$ (Kolmogorov–Smirnov test $p < 0.01$; Mann-Whitney U test $p < 0.01$), with a larger median during positive shifts of %TOC and during negative excursions in $\delta^{13}\text{C}_{\text{org}}$.

Abundance measured as number of specimens on an area of 1 dm² shows a weak positive correlation with %TOC, which seems close to, though not reaching, significance (Table 2).

Correlation between size and stratigraphic position during the T-OAE shows a two-stage pattern (displayed on Fig. 5): an initial one (between about 54 and 58 m from the base, before the maximum %TOC positive peak) of decreasing size (all samples: $r_s = -0.23$, $p < 0.001$; maximum sizes only: $r_s = -0.96$, $p = 0.003$) and a final one (between about 61 and 62 m from the base, after the %TOC positive peak) of increasing size (all samples: $r_s = 0.5117$, $p = 0.001$; maximum sizes only: $r_s = 1$, $p = 0.33$). Abundance shows an overall decreasing trend ($r_s = -0.76$, $p = 0.01$) within the T-OAE interval.

5. Discussion

5.1. Opportunistic behaviour in *Posidonotis cancellata*

Several authors have suggested opportunistic behaviour for *Posidonotis* inferred from its mode of occurrence (Hallam, 1977, p. 62, as *Pectinula*; Damborenea, 1987, p.

196; Aberhan, 1993, p. 119; Aberhan and Pálffy, 1996, p. 999). Opportunistic taxa show some particular features, such as high fecundity and short generation time, high intrinsic rates of population increase and a generalistic physiology, which allow them to rapidly increase in number in response to environmental perturbations (Levinton, 1970). Their recognition in the fossil record is not always easy, but *P. cancellata* clearly shows some of the main features of opportunistic species. It tends to appear in the logged sections as the dominant and usually the only species in thin but widespread isochronous horizons, a feature typical of opportunistic taxa (Levinton, 1970). The presence of 1–2 species of epifaunal suspension feeders at a particular horizon is comparable with the "peak of opportunist" in present-day deoxygenated areas (Caswell and Frid, 2016). During ecological crises, opportunistic species tend to appear in these monotypic associations in higher frequency (Kauffman and Erwin, 1995; Kauffman and Harries 1996, fig. 1). Although *P. cancellata* appears in the studied section at least 30 metres below the onset of the T-OAE, it is noteworthy that the higher frequencies of levels and higher abundance of specimens per level are attained during the interval recording the event itself (Fig. 5).

Fecundity and generation time are hard, if not impossible, to observe in the fossil record of most organisms. Nevertheless, fossil-association data, like the size-frequency distributions, may provide some insights on the matter (Levinton and Bambach, 1970). Taphonomic information points to autochthonous associations; no evidence of sudden burial was found on these beds, so that shell size can be related to relative time of death (Levinton and Bambach, 1970). Even though some time averaging may have occurred, *P. cancellata* beds are thin, implying short time intervals and little variations in population dynamics; unimodal samples may even represent single cohorts.

The scarcity of small shells is remarkable in the size-frequency distributions of *P. cancellata*. The smallest shell measured is about 3mm long, indicating probably the smallest measurable size; but when the distribution of the samples is analyzed (Fig. 6) it is clear that the smallest sizes, larger than that threshold, are less abundant. Taphonomic biases can be disregarded, either selective transport (both sedimentological and taphonomic evidence point to a rather quiet-water environment) or selective dissolution (dissolution affected many of the samples, but external moulds were nonetheless preserved). Present-day opportunistic bivalve species usually have the highest frequency in their smallest categories, a reflection of a high juvenile mortality rate (Levinton and Bambach, 1970). Various species studied for the same time interval in other parts of the world also show most individuals concentrated in the smaller size-classes, such as *Bositra radiata* and *Pseudomytiloides dubius* from northern England (Caswell and Coe, 2013), or *Bositra buchi* (Römer) from southwest Germany (Röhl et al., 2001). *Posidonotis cancellata* size-frequency distributions suggest that juvenile mass mortality was not particularly high in any of the populations, and mortality rates were relatively constant through time, a feature more typical of equilibrium species. Size-frequency distributions for increasing vs decreasing %TOC settings differ due to the differences in size; when the mean is subtracted from each sample, the Kolmogorov–Smirnov test shows no significant differences in distribution ($p = 0.82$). From this result, it can be concluded that only size was affected by the environmental variables analyzed here, while the reproductive strategy remained the same.

The larger median size under an increasing %TOC context (Fig. 6), together with the positive correlation of skewness with %TOC (Table 2, implying longer "positive" or right tails for length distribution in samples from environments with higher %TOC) can be related to more than one environmental variable. Percentage TOC

is one of the key characteristics used to recognize the impact of an Oceanic Anoxic Event (OAE) in the sedimentary record. The major driving force behind an OAE is supposed to have been an abrupt rise in global temperatures, probably forced by input of volcanogenic carbon dioxide, that in turn increased weathering of bedrock in subaerial and submarine environments and increased the flux of nutrients to the oceans: this would have caused an increase in organic productivity, favoring an intense oxygen demand in the water column and resulting in generalized hypoxia and increased rates of carbon burial (Jenkyns, 2010). The T-OAE, therefore, represents an interval of globally increased organic productivity during which oxygen-depleted waters characterized many marine environments, including those in the Neuquén Basin (Al-Suwaidi et al., 2010, 2016). The size increase in *Posidonotis* under increasing %TOC trends, as well as the highest abundance at the beginning of the T-OAE (Figs. 5 and 6), suggests an exploitation of plentifully available resources but, at the same time, a high tolerance to low-oxygen conditions. Although *P. cancellata* was occasionally present before oxygen levels began to decrease, its abundance increased as the rest of the benthos diminished.

Most current oxygen minimum zones (OMZ), where oxygen levels are permanently under 0.5 ml/L, show no monotonic shifts in body size for macrofauna along oxygen gradients, though in some polychaetes an increasing body size with decreasing oxygen trend has been found, which may be related to food availability (Levin, 2003). Caswell and Coe (2013) found a positive and significant regression slope of size on %TOC, similar to the results shown here, for *Pseudomytiloides dubius*. They also found an increase in size in *Bositra radiata* correlating with decreases in $\delta^{13}\text{C}_{\text{org}}$. Based on these and other responses to geochemical indicators, Caswell and Coe (2013) concluded that periods of higher primary productivity and greater oxygenation supported larger bivalves.

Hence, *Posidonotis cancellata* shows features of opportunistic species, like the high tolerance to hypoxia, its strong dominance in impoverished environments (preserved in thin extensive isochronous horizons), the strong dependence on primary productivity, but with a reproductive strategy more similar to equilibrium species (at least as regards the relatively low juvenile mortality rates).

5.2. Ecological response to the T-OAE

Macroinvertebrate faunal assemblages in the logged section are more diverse in the lower beds well below the level of the negative carbon-isotope excursion, showing a clear reduction on diversity upwards in the section. *Posidonotis cancellata* dominates (Fig. 5) the part of the section between its first record and the first T-OAE level (as identified by Al-Suwaidi et al., 2010, 2016). Apart from ammonites, only very rare *Entolium* sp., rhynchonellide and discinoid brachiopods are present in that stratigraphic interval. The latter are closely comparable to the species *Discinisca papyracea* (Goldfuss), known to occur in the lower Toarcian organic-rich shales from Swabia (Röhl et al., 2001; Schmidt-Röhl et al., 2002). At Arroyo Lapa, *Posidonotis* pavements become more abundant at the base of the T-OAE interval, intercalated with beds that in some places contain the free-living bivalve *Kolymonectes* sp. Although this last taxon may appear in dysaerobic facies in the Lower Jurassic deposits (Damborenea, 1998), it is recorded in the logged section only stratigraphically above the level of the main peak of %TOC and never forms shell-pavements. The stratigraphically highest *Posidonotis* is recorded at 63 m and the highest *Kolymonectes* at 64 m. The upward-following beds (between 64 and 69 m) show certain diversification of the fauna, containing very scarce inoceramids, lucinoids and pectinoids (*Entolium* sp.) as well as a few terebratulide brachiopods.

The abundance of *P. cancellata* reaches its highest values at the onset level of the T-OAE (Fig. 5), but it significantly decreases over the event interval up to where the taxon disappears. This stratigraphic distribution is congruent with observations on extant benthos, where densities of macrofauna are commonly depressed within the part of the OMZ where oxygen concentrations are lowest (Levin, 2003). Shell size, on the other hand, shows a two-stage response; though not strictly correlated with %TOC values in this portion of the log, it clearly shows a decreasing stratigraphic trend below the level of maximum %TOC and an increasing trend in higher levels; where %TOC reaches a maximum the species is not found. Despite surviving the conditions associated with highest %TOC, *P. cancellata* (and the whole genus) disappeared shortly thereafter, as shown by its highest occurrence in the section, in basal levels of the *Dactylioceras hoelderi* Zone (~ Serpentinum Standard Ammonite Zone).

Although the Lilliput effect (temporary body-size reduction linked to recovery events post-extinction; Twitchett, 2007; Harries and Knorr, 2009) is strictly applied for recovery events, several cases of size reduction are documented in connection with the Toarcian extinction event and also with other extinction events (e.g., Late Triassic extinction, Atkinson et al., 2019). Morten and Twitchett (2009) found a similar trend in *Pseudomytiloides dubius* from lower Toarcian sediments in northern England. That particular species apparently reached its smallest size during the late Serpentinum Zone, and disappeared shortly thereafter in the early Bifrons Zone. Fürsich et al. (2001) noticed the small size of *Parvamussium pumilum* (Lamarck) from the Toarcian of the Causses, and the same phenomenon was observed for *Bositra* and *Meleagrinella* in the T-OAE sediments of Canada (Martindale and Aberhan, 2017), and for *Bositra buchi* about the Elegans/Falciferum subzone boundary in Germany (Röhl et al., 2001). However, in other groups of organisms such as brachiopods, both decreasing and

increasing size trends were recorded prior to the early Toarcian extinction event (García Joral et al., 2018; Piazza et al., 2019) .

The overall stratigraphic size-trend observed in *P. cancellata* does not precisely correlate with any of the geochemical proxies, suggesting that environmental impoverishment should not be focused on these parameters alone, and wider oceanographic changes should be considered. Other global perturbations have been also suggested as causes of the early Toarcian extinction event. Heavily calcified organisms (*e.g.*, hermatypic corals, lithiotid bivalves, hypercalcified sponges) were selectively affected pointing to ocean acidification as another potential killing mechanism (*cf.* Trecalli et al., 2012; Hönisch et al., 2012).

Posidonotis disappeared in the section just before the minimum negative carbon-isotope values were attained (in the core of the OAE) and the genus never appeared again in the Neuquén basin. Taking into account its entire global stratigraphical range (Damborenea, 1987, 1989; Aberhan and Pálffy, 1996), the genus likely survived previous minor oceanic anoxic events, such as the Sinemurian–Pliensbachian and the Pliensbachian–Toarcian boundary events (Littler et al., 2010; Hesselbo et al., 2014; Percival et al., 2015) suggesting a relatively resilient taxon. Though *P. cancellata* survived the maximum % TOC peak of the T-OAE and even showed an increasing size trend thereafter, the abundance apparently followed a broadly decreasing pattern within the T-OAE itself.

Oxygen-poor conditions have been considered responsible for the smaller body sizes recorded in many bivalves during the Toarcian, probably due to increased juvenile mortality (Atkinson et al., 2019). Nevertheless, *P. cancellata* shows relatively low juvenile mortality and, furthermore, an increase of size that parallels %TOC. According to Levin (2003), metazoans in OMZ show some particular adaptations that allow them

to live in those environments, such as reduced blood-to-water diffusion distances, increased blood pigment affinity for oxygen, increased respiratory surfaces, and different strategies to avoid sinking in the soupy sediments typical of such environmental settings. *Posidonotis cancellata* shell morphology (and that of paper-clams in general) may be related to many of these adaptive needs; flat morphology entails a high surface/volume ratio, providing resistance to sinking, wide respiratory surfaces (mantle lobes apart from the ctenidia) with short diffusion distances. A teleplanic larval development, suggested for paper-clams, probably enabled these forms to migrate across oceanic barriers and also survive in black-shale environments (Oschmann, 1993). Metabolic adaptations cannot be identified in the fossil record, but the presence of haemoglobin in several species of bivalves belonging to different superfamilies might be invoked (Levin, 2003). If we also consider the unusual reproductive strategy (compared to other opportunistic species), it is possible that *P. cancellata* (and also the entire genus *Posidonotis*) actually represents a taxon adapted to permanent hypoxia, similar to the fauna of modern OMZs, and not to periodic hypoxia, more typical of opportunistic species. This behaviour could also explain the widest geographical distribution of the genus during the late Pliensbachian to early Toarcian time interval (Fig. 1). If that were the case, the extinction of the taxon might have been related to the environmental change driven by the end (rather than by the onset) of the T-OAE. It should be considered that the T-OAE was linked to a mass extinction event and many studies show a systematic reorganization of communities thereafter (Aberhan and Fürsich, 1997; Danise et al., 2013; Caswell and Frid, 2016; Danise et al., 2019); it is possible, therefore, that after this crisis this particular taxon could not adapt to the conditions of the new ecosystems. On the other hand, it is also possible that within the

T-OAE the oxygen level may have dropped beyond the minimum of ecological tolerance for *Posidonotis cancellata*.

6. Conclusions

The study of the size and abundance distribution of *Posidonotis cancellata* in relation with the environmental parameters determined from a section in the Neuquén Basin of Argentina shows a significantly larger shell under conditions of increasing content of total organic carbon. Abundance reaches a maximum at the beginning of the T-OAE, and then decreases till the disappearance of the species in the section. Size-frequency distribution and taphonomic information indicate that juvenile mass mortality was not particularly high in any population and mortality rates were relatively constant through time. Shell size decreases up to the level of maximum %TOC and, higher in the section, a size increase was observed to just below the extinction level of the species.

Posidonotis cancellata shows features of opportunistic species, like the high tolerance to hypoxia, the strong dominance in impoverished environments and the strong dependence on primary productivity but, on the other hand, also shows a reproductive strategy more similar to equilibrium species, with relatively low juvenile mortality rates. *Posidonotis cancellata* shows several adaptations to low-oxygen environments, like a flat morphology (providing a high surface/volume ratio, resistance to sinking, wide respiratory surfaces and short diffusion distances). The taxon may then represent a species adapted to permanent hypoxia, similar to modern faunas in oxygen minimum zones. Such ecology could explain the widespread distribution of these bivalves during latest Pliensbachian–earliest Toarcian time (interval/s of widespread hypoxia) and its extinction when environmental conditions changed. This kind of

analysis may help to improve understanding of biotic responses to large environmental disturbances under conditions of global warming, such as exist today.

Acknowledgements

This research was funded by Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET). Hesselbo and Jenkyns acknowledge funds from the Royal Society of London for a joint International Project with La Plata University. We also thank two anonymous reviewers and editor Thomas Algeo who provided helpful suggestions to improve the text. This work is a contribution to the Project IGCP-655 of the UNESCO-IUGS.

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Figure captions:

Figure 1. A- Latest Pliensbachian-earliest Toarcian global palaeogeography (modified after Cao et al., 2017); B-Location of Arroyo Lapa section in Neuquén Basin, Argentina (local palaeogeography modified after Legarreta and Uliana, 1996). Both with location of *Posidonotis* occurrences (stars).

Figure 2. **A.** Location of the Arroyo Lapa South logged section (line) (satellite image courtesy of Google Earth). **B.** General view of the studied succession at Arroyo Lapa South section (persons for scale). **C.** Detail of beds with *Posidonotis cancellata* (Leanza), coin diameter 18.2 mm.

Figure 3: Measurements of *Posidonotis cancellata* (Leanza), MLP 35885.

Abbreviations: L, length; H, height. Dashed line shows outline of measured shell-surface.

Figure 4: General aspect of slabs with different densities of specimens of *Posidonotis cancellata* (Leanza), all to the same scale. **A,** sparse, specimens separated by a distance larger than their size, MLP 35888; **B,** low density, specimens separated by distances shorter than their size, MLP 35888; **C,** high density, specimens in contact, covering almost all surface, MLP 35900; **D,** highest density overlapping valves covering entire surface, MLP 35885.

Figure 5: Logged section at Arroyo Lapa (Neuquén Basin, Argentina), showing beds with *Posidonotis cancellata*, $\delta^{13}\text{C}$ values from bulk sediment, %TOC profile, median and maximum length values, and abundance data for samples of *P. cancellata* with 5 or

more shells. General section (left, logged by Damborenea and Manceñido), and detailed section of the Pliensbachian–Toarcian boundary beds (right, logged by Al-Suwaidi et al., 2010). Local ammonite biostratigraphy by A.C. Riccardi. Solid lines in geochemical curves correspond to data used in this analysis, published by Al-Suwaidi et al. (2010); broken lines are additional data for the upper part of the section published by Al-Suwaidi et al. (2016). Grey background indicates T-OAE (*sensu* Al-Suwaidi et al., 2010, 2016).

Figure 6: Histograms showing the size (in millimetres) frequencies distribution for samples lumped under increasing and decreasing %TOC (top) and $\delta^{13}\text{C}$ (bottom) conditions.

Table 1: Main data for *Posidonotis cancellata* and geochemical variables considered for the analyses. Bold values in the first column indicates the coincidence between *P. cancellata* and geochemical samples. *dif. %TOC*= first %TOC value after the *Posidonotis* sample minus immediate previous value; *dif. $\delta^{13}\text{C}_{org}$* =first $\delta^{13}\text{C}_{org}$ value after the *Posidonotis* sample minus immediate previous value (see text).

Table 2: Results of the Spearman rank-order correlation analyses.

r_s: Spearman's coefficient; *p*: probability of non-correlation; *: significant values. *dif. %TOC*= first %TOC value after the *Posidonotis* sample minus immediate previous value; *dif. $\delta^{13}\text{C}_{org}$* =first $\delta^{13}\text{C}_{org}$ value after the *Posidonotis* sample minus immediate previous value (see text).

collect ion numbe r	dist from base (m)	N	Abund ance (ind/d m ²)	Abund ance (surf/s urf)	length (mm)					geochemical proxies				δ^{13} C	di f. δ^{13} C	%T OC	dif. %T OC
					5 maximum lengths					Med ian	Vari ance	Skew ness	Kurt osis				
MLP35 870	24. 55	1 1	7.2	0.14	17 .5	17 .1	16 .8	16 .6	16 .6	12.0	26.1	-0.30	- 1.82	- 27. 45	- 0. 64	0. 34	- 1.7 1
MLP35 871	33. 1	1 4	12.4	0.24	20 .8	18 .9	17 .5	17 .5	16 .6	14.9	9.8	0.23	- 0.88	- 24. 73	- 0. 59	2. 28	1.6 7
MLP35 873	36. 05	5	7.9	0.20	17 .5	13 .6	13 .0	12 .4	10 .1	13.0	7.1	0.81	1.94	- 26. 24	0. 55	3. 66	2.8 4
MLP35 874- MLP35 876	36. 35	6 0	29.2	0.40	24 .6	23 .0	22 .8	20 .6	19 .7	11.7	22.1	0.48	- 0.44	- 26. 63	- 0. 39	4. 07	0.4 0
MLP35 878	39. 05	2 1	18.3	0.34	17 .8	16 .8	16 .0	15 .1	14 .7	12.5	9.4	-0.03	- 1.13	- 26. 91	- 1. 04	3. 20	- 0.0 9
MLP35 879	42. 45	2 5	40.2	0.44	20 .3	19 .2	18 .7	15 .3	13 .4	9.6	19.9	0.84	0.12	- 26. 65	- 2. 43	4. 25	2.0 2
MLP35 881	42. 85	6	24.5	0.52	10 .9	10 .9	10 .7	8. 7	7. 3	9.7	3.5	-0.36	- 2.49	- 25. 53	1. 12	3. 62	- 0.6 3
MLP35 882	43. 05	8	26.0	0.28	14 .8	13 .7	10 .9	10 .2	9. 7	10.0	10.1	0.17	- 0.55	- 25. 53	1. 12	3. 62	- 0.6 3
MLP35 883- MLP34 884	46. 05	2 7	11.0	0.16	16 .7	16 .6	14 .4	13 .7	13 .4	10.2	11.1	0.04	- 0.41	- 24. 30	0. 29	3. 05	- 0.8 4
MLP35 885	54. 55	1 6 8	51.4	0.62	24 .3	21 .0	20 .9	19 .8	19 .1	13.3	13.1	0.08	- 0.42	- 26. 78	- 4. 89	2. 63	0.5 5
MLP35 887- MLP35 888	54. 65	8 4	63.4	0.68	20 .7	20 .7	19 .3	19 .3	17 .9	12.0	12.3	0.30	0.18	- 26. 78	0. 47	2. 63	1.6 8
MLP35 886	55. 55	7	64.3	0.71	17 .1	16 .5	13 .9	13 .5	13 .3	13.5	5.6	0.05	- 0.38	- 26. 81	- 0. 50	4. 88	0.5 7
MLP35 890	55. 85	1 5	56.6	0.52	18 .2	14 .1	13 .1	12 .2	11 .1	10.6	9.1	1.09	1.69	- 25. 43	1. 38	4. 06	- 0.8 2
MLP35 892	56. 65	7	22.0	0.36	15 .6	15 .5	14 .5	13 .5	12 .1	13.5	9.7	-1.45	2.34	- 25. 87	0. 43	2. 73	- 0.9 8

MLP35 893	57. 3	1 2	34.7	0.88	15 .1	14 .0	13 .3	13 .1	12 .3	11.6	4.6	-0.46	0.42	- 25. 44	- 0. 46	1. 75	0.3 1
MLP35 894	57. 65	5	23.2	0.24	12 .4	9. 6	8. 7	8. 3	5. 3	8.7	6.5	0.00	1.37	- 25. 90	- 0. 61	2. 06	2.3 5
MLP35 895	61. 05	2 2	27.8	0.39	17 .6	15 .5	12 .2	12 .0	12 .0	10.0	7.8	1.20	1.64	- 27. 88	- 1. 00	4. 36	- 1.9 7
MLP35 896	61. 55	8	25.9	0.29	18 .8	15 .5	15 .0	13 .3	12 .5	12.9	12.9	-0.22	0.88	- 27. 91	- 0. 03	5.6 7	1.3 1
MLP35 897	61. 85	7	6.2	0.14	20 .7	20 .6	16 .9	15 .8	14 .1	15.8	23.1	-0.32	- 1.11	- 28. 48	- 0. 57	0.6 2	- 5.0 5

Table 1 - Main data for *Posidonotis cancellata* and geochemical variables considered for the analyses. Bold values in the second column indicates the coincidence between *P. cancellata* and geochemical samples. **dif. %TOC**= first %TOC value after the Posidonotis sample minus immediate previous value; **dif. $\delta^{13}\text{C}$** =first $\delta^{13}\text{C}$ value after the Posidonotis sample minus immediate previous value (see text).

		$\delta^{13}\text{C}$	<i>dif.</i> $\delta^{13}\text{C}$	%TOC	<i>dif.</i> %TOC
max. length	r_s	-0.36	-0.36	0.06	0.14
	p	0.13	0.13	0.79	0.56
5 max. length	r_s	-0.28	-0.36	-0.12	0.06
	p	*0.01	*0.00	0.25	0.57
median	r_s	-0.28	-0.08	-0.21	-0.06
	p	0.25	0.73	0.39	0.80
variance	r_s	-0.40	-0.30	-0.17	-0.16
	p	0.09	0.22	0.49	0.50
skewness	r_s	0.04	-0.02	0.50	0.31
	p	0.86	0.92	*0.03	0.20
kurtosis	r_s	0.12	0.23	0.31	0.24
	p	0.63	0.34	0.19	0.31
Abundance (ind/dm²)	r_s	0.03	0.05	0.45	0.29
	p	0.90	0.84	0.06	0.23
Abundance (surf/surf)	r_s	0.09	0.06	0.31	0.22
	p	0.71	0.82	0.20	0.37

Table 2 - Results of the Spearman rank-order correlation analyses.

r_s : Spearman's coefficient; p : probability of non-correlation; *: significant values.

dif. %TOC= first %TOC value after the Posidonotis sample minus immediate previous value; **dif. $\delta^{13}\text{C}$** =first $\delta^{13}\text{C}$ value after the Posidonotis sample minus immediate previous value (see text).

HIGHLIGHTS

- *P. cancellata* size and abundance data from Toarcian relate to OAE geochemical proxies
- Shell size correlates positively with %TOC, probably linked to primary productivity
- Maximum abundance at the onset of the T-OAE, and then decreasing until its extinction
- *P. cancellata* shows opportunistic species features, but with a low juvenile mortality

P. cancellata anatomical features suggest adaptation to dysaerobic environments.

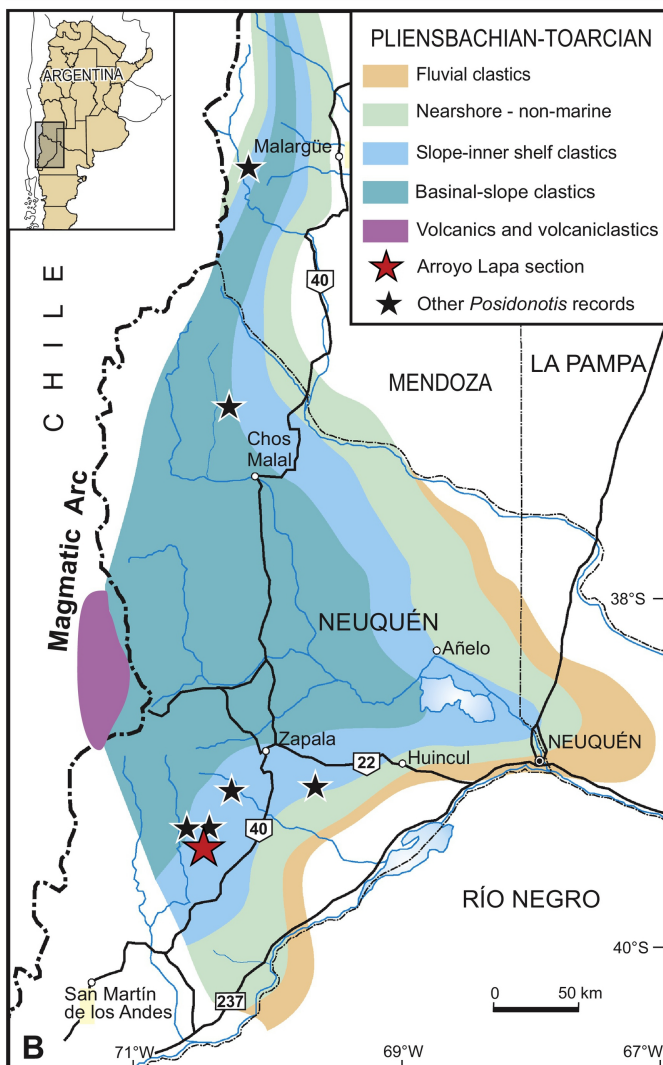
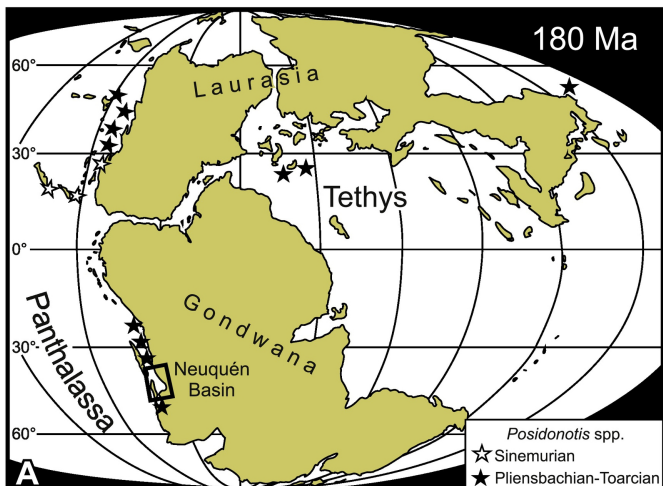


Figure 1



Figure 2

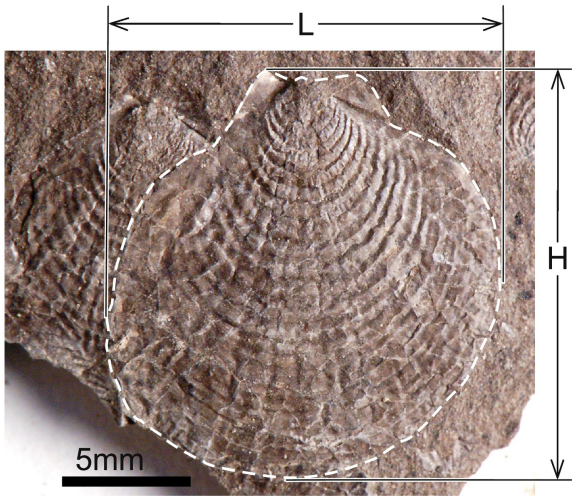


Figure 3



Figure 4

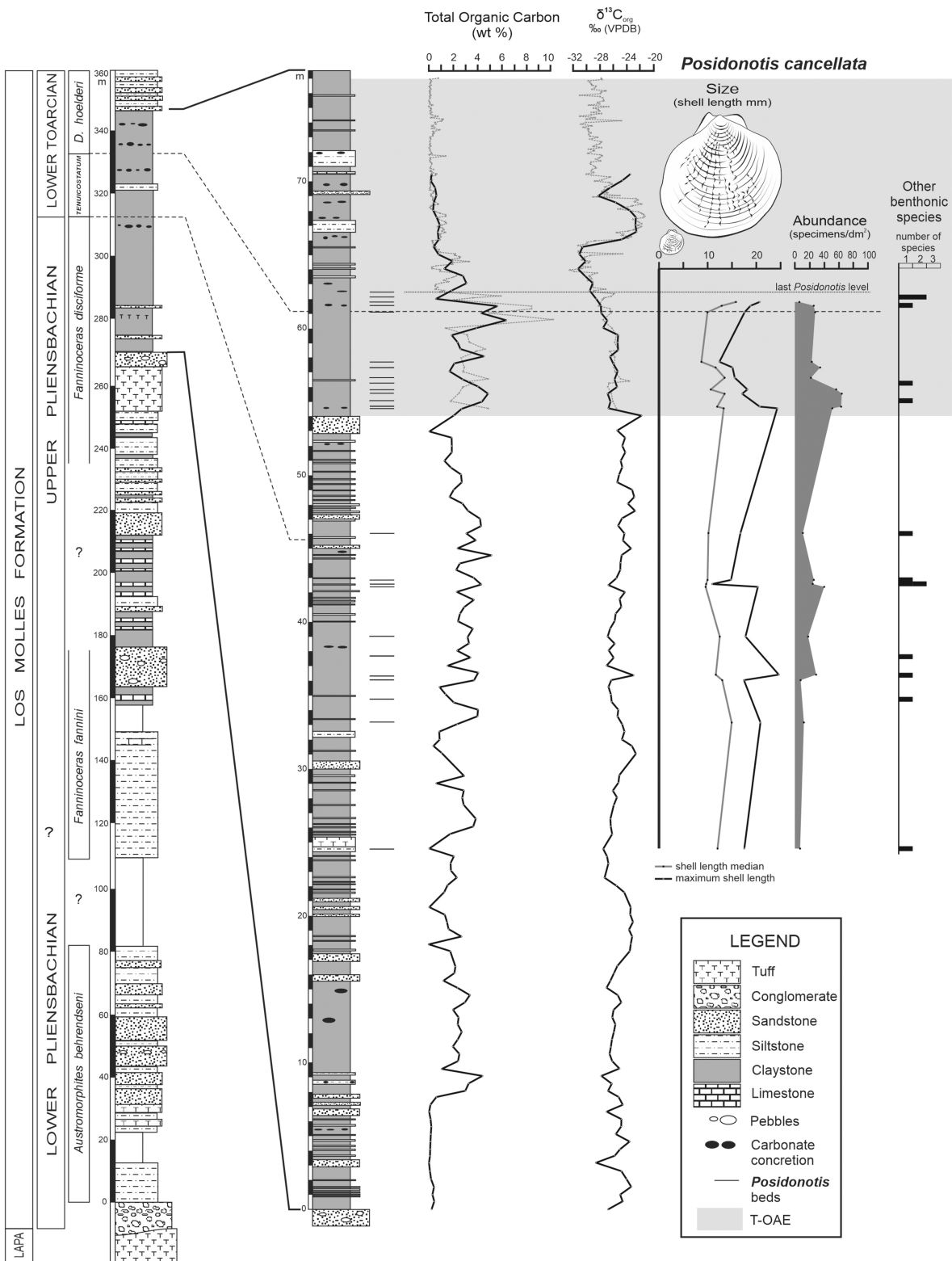


Figure 5

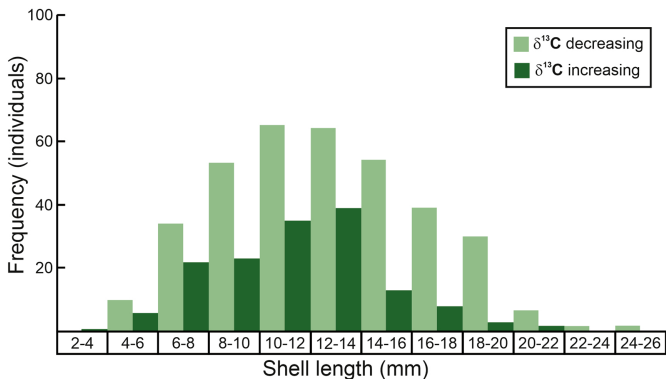
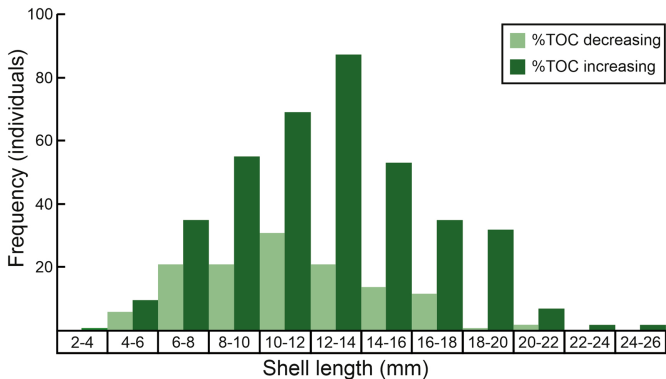


Figure 6